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## Crystal and Molecular Structure of (CH<sub>3</sub>)<sub>2</sub>Sn[C<sub>6</sub>H<sub>5</sub>Cr(CO)<sub>3</sub>]<sub>2</sub>

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TECHNICAL REPORT NO. 28

#### Summary

1. 1. 1. 6-

 $n^4$ ,  $n^4$ -Dimethyldiphenylstannane-bis-tricarbonylchromium,  $(CH_3)_2Sn[C_6H_5Cr(CO)_3]_2$  crystallizes in the orthorhombic space group Pna2<sub>1</sub> with Z = 4, a = 2663(2), b = 1032.0(8), c = 762.7(3) pm, V = 2096 x 10<sup>6</sup> pm<sup>3</sup>. The crystal structure has been determined by the heavy atom method and refined to final R values of R = 0.055 and R = 0.059 for 1789 independent reflections using  $M_0$ - $K_{\alpha}$  radiation. In the lattice, the molecule is disordered in that one of the  $Cr(CO)_3$  groups can be on either side of the phenyl ring to which it is bonded; the unit cell contains molecules of both configurations in equal numbers. The geometry of the four groups around the tin atom is approximately tetrahedral; thus, anomalous spectral properties of this compound cannot be attributed to molecular distortions of the normal tin geometry.

### Introduction

Several years ago, Willeford, Zuckerman and coworkers reported the results of a spectroscopic investigation of several arene metal carbonyl complexes of tin-containing ligands [2]. All but one of the complexes studied showed tin-methyl stretching absorptions in the infrared practically identical with those of the uncomplexed ligands. That one exception was  $\eta^6$ ,  $\eta^6$ -dimethyldiphenyl-stannane-bis-tricarbonylchromium,  $(CH_3)_2Sn[C_6H_3Cr(CO)_3]_2$ . This complex showed no absorption in the infrared region where the symmetric tin-methyl stretching

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vibration would be expected to occur. Raman spectroscopy revealed a strong polarized band coincident with the infrared absorption assigned to the asymmetric tin-methyl vibration. Thus, the symmetric and asymmetric tin-methyl vibrations in this complex apparently are degenerate. Gross distortion of the tin atom in this complex from its usual tetrahedral symmetry was suggested as a possible explanation for this unusual observation. A crystallographic investigation was undertaken to determine the validity of this conjecture.

### Experimental

First attempts to grow single crystals of the complex resulted only in twinned crystals. Satisfactory crystals were finally obtained by careful layering of oxygen-free hexane over a concentrated solution of the complex in benzene. After several days standing at ambient temperature, crystals formed at the benzene-hexane interface which were collected, washed with hexane, and dried under vacuum. A crystal, approximately 0.4 x 0.4 x 0.8 mm. was mounted on a Syntex P2, four circle automatic diffractometer. Lattice parameters (see Table 1) were determined by least squares from the setting angles of 15 reflections from different parts of the reciprocal space. Mo-K radiation was used (graphite monochromator,  $\lambda$  = 71.069 pm) for intensity data collection ( $\omega$ -scan,  $\Delta \omega$  = 0.9°, 1789 independent reflections,  $2^{\circ} \stackrel{\leq}{=} 29 \stackrel{\leq}{=} 46^{\circ}$ ). The data were corrected for Lorentz and polarization factors, but not for absorption. The structure was solved by the heavy atom method and refined by full matrix least squares. Only the metal atoms were treated anisotropically. The hydrogen atom parameters were kept constant during refinement. For 1573 structure factors with  $F_0 \ge 3.92 \sigma(F_0)$  final R = 0.055 and  $R_{w} = 0.059$  were obtained  $(1/W = \sigma^2 (F_0))$ .

Table 2 shows the final atomic parameters, while Table 3 gives intramolecular distances and angles.

### Description of the structure and discussion

For this compound two favorable isomers can be imagined:

For steric reasons, "in/in" complexes are formed only if there is a metal-metal bond [as in Fe<sub>2</sub>(CO)<sub>4</sub>( $\pi$ -C<sub>5</sub>H<sub>5</sub>-SiMe<sub>2</sub>- $\pi$ -C<sub>5</sub>H<sub>5</sub>) [3]]. Rotation of the phenyl rings about the Sn-C<sub>phenyl</sub> axes is hindered by interactions between the phenyl rings and is not to be expected.

Because of the disorder of one of the  $Cr(CO)_3$  moieties, both "out/out" and "in/out" isomers are present in the crystal. The figure shows the superposition of both isomers. Whereas the positions of the  $Ph_2SnMe_2$  moiety and one of the  $Cr(CO)_3$  groups [the one which is "out" in both isomers, Cr(1)] are not affected by the disorder, the positions of Cr(2) and Cr(3) are only half-occupied. The CO ligands of the disordered  $Cr(CO)_3$  moiety are positioned in such a way that two oxygen atoms  $\{O(4) \text{ and } O(5)\}$  and one carbon atom  $\{C(6)\}$  on Cr(2) have common coordinates with the analogous atoms at Cr(3) of the neighboring molecule  $\{O.5-x, -0.5+y, 0.5+z\}$ .

Because of the disorder in the crystal, the  $Cr-C_{CO}$  and  $Cr-C_{ph}$  distances and the Cr-C-O angles are found to differ considerably and have large standard deviations (see Table 3). However, their mean values are quite similar. The

CO ligands and the carbons of the phenyl groups are staggered. The phenyl rings are planar within 1.8 standard deviations. Whereas Cr(1) is located above the center of the phenyl ring, the distances of Cr(2) and Cr(3) to C(21) and C(22) are somewhat longer than to C(24) and C(25) (see Table 3). The Cr-phenyl plane distances are 170.1 (Cr 1), 162.6 (Cr 2) and 163.8 (Cr 3) pm respectively. Because of the disorder, a more detailed discussion of the bond lengths and angles is not justified.

The tin atom is located 8 and 9 pm outside the best planes of the phenyl rings (towards the "out" position). Its coordination is approximately tetrahedral. The Sn-C<sub>phenyl</sub> bond lengths (218 pm) are slightly longer than in tetraphenyltin [4] or in triphenyltin compounds [5-8], where bond lengths of 212-216 pm are found.

The results give a dimethyltin carbon-tin-carbon angle of 115.5(7)°, opened from the tetrahedral value at the expense of the diphenyltin angle of 105.7(5)° which is somewhat closed. Structural data on the free dimethyldiphenylstannane ligand, or on any simple, mixed methylphenyltin system are lacking [8]. Apparently there are structural data for only two compounds in which both unsubstituted methyl and phenyl groups are attached to the same tin atom [9] (the structures of Ph<sub>3</sub>SnCH<sub>2</sub>I [10] and a ring system based upon 1,8-bis-dimethylstannylnaphthalene [11] do not offer meaningful comparison). These compounds, 4-chloro- and 4-bromo-1,2,3,4-tetraphenylbutadienyldimethylphenylstannane, exhibit closed dimethyltin carbon-tin-carbon angles of 105.5° and 104.5°, respectively, while the mean methylphenyltin angles are 107.7° and 108.4°, respectively. The bonds involving the vinylic system are opened to even larger angles [9]. This is opposite to what is observed in the title compound, where the relative magnitudes of the dimethyltin and diphenyltin angles (the methylphenyltin angles are intermediate in value) reflect a greater concentration of g-character in the bonds the tin atom exerts to hold the methyl groups, and a greater concentration of p-character toward the phenyl groups. Thus the phenyl groups in this case are exhibiting a greater electronegativity.

The direction of the relative opening of the dimethyltin angles is in accord with the suggestion of Willeford and Zuckerman [1], but a distortion of this small magnitude (6.0° from tetrahedral, 64.5° from linear), while of interest, obviously cannot lie at the root of the absence of the symmetric Sn-C stretching absorption in the infrared spectrum and the presence in the Raman spectrum of only a single absorption in this region. Other rationalizations of these spectroscopic data must be sought.

### Acknowledgments

We thank Prof. Dr. E. O. Fischer for his interest and for making available the facilities of the Institute. B.R.W. thanks the Deutscher Akademischer Austauschdienst, Bonn, for financial support and the Fulbright Kommission, Bonn, for a travel grant.

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# Figure Caption

A perspective drawing of the molecular structure of  $\eta^6, \eta^6\text{-dimethyldiphenylstannane-bis-tricarbonylchromium,}$   $(CH_3)_2 Sn[C_6H_5Cr(CO)_3]_2. \quad \text{The disordered part of the}$  molecule is drawn with light dotted lines. The hydrogen atoms are omitted for clarity.

Table 1

Crystal Data

Formula (molecular weight)  $C_{20}H_{16}Cr_{2}O_{6}Sn$  (575.0)

Space group  $Pna2_1 (Z = 4)$ 

Cell constants: a,b,c 2663(2), 1032.0(8), 762.7(3) [pm]

V 2096•10<sup>6</sup> [pm<sup>3</sup>]

Density (calc.) 1.82 [g/cm<sup>3</sup>]

Linear absorption coeff. (Mo-K $_{\alpha}$ ) 22.8 [cm $^{-1}$ ]

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Tab

Atom	x/a	e/k	2/2	ø	Atom	ж/в	3/6	2/2	m
Sn	0.13242(3)	0.01223(8)	0.1		Cr3	0.2600(2)	-0.1047(4)	0.3246(6)	
Cr1	0.03568(7)	-0.2736(2)	0.1423(4)		Cr2	0.2634(2)	0.1526(4)	0.0753(8)	
	0.1011(7)	0.095(2)	0.337(3)	4.8(4)	C02	0.1156(7)	0.116(2)	-0.140(3)	4.7(4)
<b>C10</b>	0.1087(5)	-0-187(1)	0.072(2)	3.0(3)	020	0.2175(4)	0.005(1)	0.139(2)	2.9(3)
C11	0.0844(6)	-0.230(2)	-0.087(2)	4.2(3)	C21	0.2346(7)	0.069(2)	0.281(3)	5.5(4)
C12	0.0665(7)	-0.359(2)	-0.095(3)	5.2(4)	, C22	0.2889(7)	0.074(2)	0.299(3)	5.5(4)
513	0.0753(7)	-0.445(2)	0.042(2)	4.5(4)	C23	0.3165(6)	0.006(2)	0.189(2)	3.8(3)
C14	0.0988(7)	-0.408(2)	0.184(3)	5.3(4)	C24	0.2963(6)	-0.062(2)	0.053(2)	4.2(4)
C15	0.1164(6)	-0.276(1)	0.201(2)	3.9(3)	<b>C25</b>	0.2445(6)	-0.062(2)	0.028(2)	3.8(3)
5	0.0200(7)	-0.281(2)	0.368(3)	4.7(4)	ð	0.1765(4)	0.342(1)	0.120(2)	5.9(3)
6	0.0081(5)	-0.287(1)	0.514(2)	7.3(4)	C41	0.218(1)	0.278(3)	(9)601.0	5.0(7)
S	0*0255(5)	-0.345(1)	0.094(3)	3.9(3)	C42	0.1970(9)	0.361(2)	0.005(4)	1.8(4)
05	-0.0647(4)	-0.389(1)	0.062(2)	5.1(3)	90	0.3275(5)	0.372(1)	0.095(3)	7.6(3)
C3	0.0068(5)	-0.114(1)	0.123(3)	3.9(3)	C51	0.307(1)	0.282(3)	0.092(6)	4.5(6)
60	-0.0097(4)	-0.010(1)	0.127(2)	4.8(2)	c>5	0.287(1)	0.382(5)	-0.014(4)	2.9(6)
H11	0.081	-0-174	-0.187	5.0	90	0.255(1)	0.195(3)	-0.191(5)	9.3(8)
H12	0.047	-0.387	-0.194	5.0	061	0.2513(9)	0.111(3)	-0.266(4)	5.2(6)
H13	450.0	-0.533	0.029	5.0	062	0.251(1)	0.195(3)	-0.312(5)	7.3(8)
H14	0.104	-0-471	0.277	5.0	H23	0.353	900.0	0.208	2.0
H15	0.135	-0.252	905.0	5.0	H24	0.319	-0.104	-0.028	5.0
H21	0.213	0.115	0.354	6.0	1125	0.250	-0.108	-0.008	0.3

table 2 continued

atom	· B <sub>11</sub>	B <sub>22</sub>	B <sub>33</sub>	B <sub>12</sub>	B <sub>13</sub>	B <sub>23</sub>
Sn	2.24(3)	3.64(4)	4.18(4)	-0.17(3)	0.03(6)	-0.11(7)
Cr1	2.24(8)	2.76(9)	3.94(13)	-0.21(7)	-0.15(9)	0.13(10)
Cr2	2.7(2)	3.4(2)	3.5(3)	-0.3(2)	-0.1(2)	0.0(2)
Cr3	1.6(2)	2.1(2)	2.2(2)	0.1(1)	-0.2(2)	-0.1(2)

The anisotropic thermal parameter is defined:

$$T = \exp[-1/4(h^2a^{*2}B_{11} + k^2b^{*2}B_{22} + 1^2c^{*2}B_{33} + 2hka^*b^*B_{12} + 2hla^*c^*B_{13} + 2klb^*c^*B_{23})]. B_{ij} in 10^4pm^2.$$

Table 3: Di	stances (in	pm) an	d angles	(in °):		
5n-C01	216(2)			Sn-CO2	217(2)	
Sn-C10	218(1)			Sn-C20	218(1)	
•				Cr1-C10	221(1)}	
Cr1-C1	177(2)	mean:	181	Cr1-C11	222(2)	
Cr1-C2	102(1)	mean:	101	Cr1-C12	217(2)	
Cr1-C3	183(1)	•		Cr1-C13	219(2)	mean: 220
C1-01	116(3)			Cr1-C14	220(2)	
C2-02	117(2)				220(2)	
C3-03	116(2)			Cr1-C15		
Cr2-C41	179(3)			Cr2-C20	208(1)	
Cr2-C51	178(3)	mean:	189	Cr2-C21	195(2)	
Cr2-C6	210(4)			Cr2-C22	201(2)	meam: 216
C41-04	129(3)			Cr2-C23	225(2)	
<b>C51-O</b> 5	107(3)			Cr2-C24	239(2)	
C6-062	92(5)			Cr2-C25	230(2)	
Cr3-C42	183(3)		•	Cr3-C20	220(1)	
Cr3-C52	177(3)	mean:	190	Cr3-C21	195(2)	
	210(3)			Cr3-C22	201(2)	mean: 216
Cr3-C6*	105(3)			Cr3-C23	216(2)	mean; 2.0
C42-04*				Cr3-C24	233(2)	
c52-05° ce-061	136(3) 105(4)			Cr3-C25	234(2)	
00-00 (				<b>C1-Cr1-C</b> 2	88.2(8	1)
CO1-Sn-CO						
CO1-Sn-C1				C1-Cr1-C3		
CO1-Sn-C2	20 106.4(6)	)		C2-Cr1-C3		
CO2-Sn-C1	10 109.0(6)	)		C41-Cr2-C	-	
CO2-Sn-C2	20 109.7(6)	)			26	
C10-Sn-C	20 105.7(5)	)		C51-Cr2-C		
				C42-Cr3-		
• 0.5-x.	-0.5+y, 0.5	+2		<b>C6-Cr3-</b> C		
				<b>C6-C</b> r3-C	42 88.2(	12)



